The axial transsacral approach to interbody fusion at L5–S1

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Lumbosacral interbody fusion may be indicated to treat degenerative disc disease at L5–S1, instability or spondylolisthesis at that level, and severe neural foraminal stenosis resulting from loss of disc space height. In addition, L5–S1 interbody fusion may provide anterior support to a long posterior fusion construct and help offset the stresses experienced by the distal-most screws. There are 3 well-established techniques for L5–S1 interbody fusion: anterior lumbar interbody fusion, posterior lumbar interbody fusion, and transforaminal lumbar interbody fusion. Each of these has advantages and pitfalls. A more recently described axial transsacral technique, utilizing the presacral corridor, may represent a minimally invasive approach to obtaining lumbosacral interbody arthrodesis. Biomechanical studies demonstrate that the stiffness of the axial rod is comparable to existing fixation devices, suggesting that, biomechanically, it may be a good implant for obtaining lumbosacral interbody fusion. Clinical studies have demonstrated good early results with the use of the axial transsacral approach in obtaining lumbosacral interbody fusion for degenerative disc disease, spondylolisthesis, and below long posterior fusion constructs. The technique is exacting and complications can be major, including rectal perforation and fistula, loss of correction, and pseudarthrosis.

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Interbody fusion at L5–S1 is indicated for specific pathology including degenerative disc disease, instability, neural foraminal stenosis, and protection of S-1 screws at the end of long fusion constructs to the sacrum. Interbody fusion at L5–S1 may be achieved anteriorly (anterior lumbar interbody fusion [ALIF]), posteriorly (posterior lumbar interbody fusion [PLIF]), or via the neural foramen (transforaminal lumbar interbody fusion [TLIF]). These techniques have specific advantages and pitfalls, making none of them the ideal approach for lumbosacral fusion. The more recently described transsacral axial lumbar interbody fusion (AxiaLIF) addresses some of the concerns of the previous fusion techniques. Early evidence suggests that the technique may be performed safely to achieve L5–S1 fusion for degenerative disc disease, spondylolisthesis, and at the distal end of long posterior fusion constructs. This review will focus on the biomechanical studies, early clinical outcomes, and complications of the axial transsacral approach to lumbosacral interbody fusion.

**Current Approaches to L5–S1 Interbody Arthrodesis**

There are 3 well-established methods to obtain L5–S1 interbody arthrodesis. An ALIF requires a retroperitoneal approach to reach the L5–S1 interspace below the bifurcation of the aorta into the left and right common iliac arteries. This direct access allows for release of the anterior longitudinal ligament and restoration of sagittal alignment using a lordotic graft with a large footprint. The anterior approach can help restore sagittal balance, increase neural foraminal height, and lead to high fusion rates. However, the exposure risks injury to the iliac vessels (especially the thin-walled common iliac vein that is adjacent to the L5–S1 disc space on the left side) and the sympathetic plexus, which can cause retrograde ejaculation in males. The PLIF and TLIF cages are smaller than ALIF cages, necessitated by the limited space for entry around the cauda equina and nerve roots; the smaller size results in a smaller footprint and a potentially lower rate of arthrodesis. Retraction to place these cages can result in neurological injury. In the majority of cases, the direct lateral interbody approach (DLIF or XLIF [extreme-lateral interbody fusion]) cannot be used to obtain interbody arthrodesis at L5–S1 because of the overhang...
of the iliac wing and obstruction by the nerve roots of the lumbosacral plexus.27

The Axial Transsacral Approach to L5–S1 Interbody Arthrodesis

The axial transsacral approach to interbody fusion at L5–S1 allows for placement of a cylindrical cage (AxiaLIF, TranS1) through the presacral corridor. The AxiaLIF implant has a reverse thread pitch to provide interspace distraction during implantation (Fig. 1). This technique has certain potential advantages including a muscle-sparing approach and complete preservation of the anulus fibrosus.22,28 The minimally vascular presacral corridor allows for safe implant placement with little risk for vascular injury.22 However, the trajectory must be very precise as the implant is navigated posterior to the rectum, and rectal injury has been reported in conjunction with this technique.4,10 Biomechanical and preliminary clinical data suggest good short-term outcomes for axial lumbar interbody fusion for degenerative disc disease, spondylolisthesis and as anterior support caudal to long fusion constructs.5,9,17,20

Biomechanical Evaluation

Biomechanical studies have shown that the axial transsacral introduction of a fusion cage is technically feasible and that the fixation properties of this technique are comparable to those of existing implants for lumbosacral arthrodesis.6,9,17,20 Ledet and colleagues27 mechanically tested 24 bovine lumbar motion segments in sagittal and lateral bending, torsion, and axial compression after drilling an axial canal and implanting a fixation rod within the drilled canal. Drilling had little effect on stiffness and range of motion of the specimens. However, specimens implanted with the axial rod demonstrated significant decreases in range of motion and increases in stiffness relative to the intact state. Compared with results reported for existing anterior, posterior, and interbody instrumentation, specifically femoral ring allograft, BAK cages, Brantigan ALIF and TLIF implants, Harms cages, and Kaneda, Isola, and University plating systems, lateral and sagittal bending stiffness of the axial rod was greater, whereas stiffness in extension and axial compression was similar to plate and rod constructs. Torsional stiffness was similar to that of interbody constructs and lower than that of plate and rod constructs. Thus the stiffness of the axial rod is comparable to or greater than that of existing fixation devices, which suggests that biomechanically it is a good implant for obtaining lumbosacral interbody fusion.17

Erkan and colleagues8 have tested 2-level fusion (L4–S1) using the axial rod technique. Six human cadaveric L4–S1 motion segments were tested in axial torsion, lateral bending, and flexion-extension following intact, stand-alone AxiaLIF (2-level rod), and AxiaLIF7 with posterior fixation and either facet screw or pedicle screw placement. At the L4–5 level in axial torsion and flexion-extension, none of the surgical treatments showed any statistical significance. In lateral bending, the posterior fixation devices had significantly higher construct stability compared with the stand-alone AxiaLIF. At the L5–S1 level in axial torsion and lateral bending, none of the surgical treatments showed statistically significant differences. However, in flexion-extension, the stand-alone AxiaLIF had significantly greater range of motion than the posterior fixation techniques, suggesting that AxiaLIF should be performed in conjunction with posterior fixation to achieve greater stability for a successful arthrodesis.9

Surgical Procedure

The patient undergoes a standard bowel preparation 24 hours prior to surgery. The procedure is performed with the patient prone and a pillow placed beneath the pelvis for elevation. Preoperative antibiotic agents are administered. The sacrococcygeal region is prepared and draped in the standard sterile fashion. If a prior posterior spinal fusion had been performed at the same operative setting, the posterior wound is closed and the patient is re-prepped and draped for the axial interbody procedure. This re-prepping minimizes infection risk and allows for optimal patient positioning in lumbosacral lordosis to help with implant targeting into the vertebral body of L-5. A 3-cm transverse incision is made at the right side of the coccyx and carried through the subcutaneous tissue to the fascia. The anterior surface of the coccyx is located with a curved Kelly clamp. Using biplanar fluoroscopy, a blunt dissecting tool (TranS1) is then advanced along the anterior face of the sacrum to the S1–2 level where it is docked (Fig. 2A). The inner blunt stylet is exchanged for a guide pin, which is introduced into the S-1 vertebral body, across the L5–S1 disc space under fluoroscopic visualization (Fig. 2B). A series of dilators opens the osseous path, allowing for placement of a working cannula; through this cannula, a path is drilled from the anterior sacrum through S-1 into the L5–S1 disc space. Radial cutting instruments are used to perform the discectomy and prepare the endplates for fusion (Fig. 2C). Cutting instruments should not be turned 360° as they may invade

P. S. Issack, S. Y. Kotwal, and O. Boachie-Adjei
the spinal canal. After discectomy, the disc space is filled with bone graft material composed of local bone removed from the vertebral body and demineralized bone matrix. The appropriate length AxiaLIF implant is inserted over a guide pin through a protective cannula, transfixing the L5–S1 level (Fig. 2D). The wound is irrigated and closed after removal of guide wires and cannulas.

Li and colleagues,18 in a human cadaveric study, dissected the presacral space in 16 pelvic specimens and observed the position of the entry guide pin for AxiaLIF in relation to the pelvic splanchnic nerves. The pelvic splanchnic nerves limited the dissection of the lower rectum, and the minimum distance from the guide pin to the pelvic splanchnic nerves was as small as 4 mm. Clearly the margin of error is small, and the accurate placement of the blunt stylet through which the guide pin is passed is the most critical step in the operation.

Treating Degenerative Disc Disease and Spondylolisthesis With AxiaLIF

The majority of clinical studies on the axial interbody technique involve treatment of degenerative disc disease and spondylolisthesis.3,11,12,22,30,31 Aryan and colleagues3 reviewed data obtained in 35 patients with back pain due to lumbar degenerative disc disease, degenerative scoliosis, or isthmic spondylolisthesis treated with AxiaLIF and recombinant human bone morphogenetic protein. Twenty-three patients underwent supplemental posterior pedicle screw fixation. In 32 patients (91%), there was clinical and radiographic evidence of L5–S1 interbody fusion at a mean follow-up of 17.5 months. Tobler and Ferrara30 prospectively followed up 26 patients with degenerative disc disease who underwent AxiaLIF with posterior pedicle screw fixation for 2 years. The fusion was at 1 level (L5–S1) in 17 patients and at 2 levels (L4–S1) in 9 patients. Interbody fusion was achieved at 1 year in 22 patients and 2 years in 23 patients. One patient with a pseudarthrosis underwent successful revision posterolateral fusion. In a larger retrospective study, Tobler and colleagues31 evaluated 156 patients who underwent an L5–S1 interbody fusion in which an AxiaLIF rod was used (Fig. 3). There were significant improvements in pain and mean Oswestry Disability Index scores at the 2-year follow-up. Radiographic evidence of interbody fusion was observed in 94% of the patients (145 of 155).

Gerszten and colleagues11 reported on 26 patients with Grade 1 or Grade 2 symptomatic L5–S1 isthmic spondylolisthesis who underwent L5–S1 AxiaLIF and posterior pedicle screw fixation. Approximately half the patients showed a reduction of at least one grade. Axial pain improved after AxiaLIF with a 66% reduction from baseline. The fusion rate at 2 years was 100% with outcome in 81% of patients deemed excellent or good according to Odom criteria.11

AxiaLIF Caudal to Long Fusion Constructs

In adult deformity surgery, long posterior fusion constructs must often be extended to include L5–S1 in situations where there is lumbosacral disc degeneration,
spondylosis, degenerative spondylolisthesis, or a fractional lumbosacral curve greater than 15° in magnitude.\textsuperscript{1,16,24} To protect distal fixation, particularly S-1 screws, the addition of iliac screws and anterior interbody support at L5–S1 is recommended to offset load on distal fixation.\textsuperscript{3,8,34} While interbody fusion at L5–S1 caudal to long fusion constructs has traditionally been achieved using an ALIF, PLIF, or TLIF, the axial interbody approach has recently been demonstrated to provide anterior support distal to long posterior fusions at short-term follow-up (Fig. 4).

Anand and colleagues\textsuperscript{1} evaluated circumferential fusion for lumbar degenerative scoliosis in 12 patients, of whom 5 had fusions to L5–S1 performed using the axial technique. There were no complications. Blood loss was less than 200 ml. Fusions extended as proximal as T-12 in these 5 AxiaLIF cases.\textsuperscript{1} In a retrospective study of 28 patients who underwent minimally invasive correction and fusion over 3 or more levels for adult scoliosis, Anand and associates\textsuperscript{2} performed 13 one-level (L5–S1) and 4 two-level (L4-S1) axial interbody fusions. At a mean follow-up of 22 months, fusion had been achieved in all patients. There were no complications related to the AxiaLIF procedure. Blood loss was 231 ml (including that associated with the placement of posterior instrumentation). These results suggest good outcomes using the axial transsacral technique to fuse L5–S1 below posterior fusion constructs extending up to T-12.\textsuperscript{1,2}

Issack and Boachie-Adjei retrospectively examined 9 patients who underwent axial interbody fixation and fusion caudal to long fusion constructs for adult scoliosis. There were 4 one-level and 5 two-level procedures. Fusions in this series were longer than those described above, with 6 patients having fusions extending proximally to T-10 or higher (up to T-3). There were 2 pseudarthroses, and no major complications occurred. There were significant improvements in the pain, self-image and satisfaction with management domains of the SRS-22 (Scoliosis Research Society questionnaire).\textsuperscript{15}

\textbf{Fig. 3.} \textbf{Upper:} Anteroposterior and lateral radiographs demonstrating L5–S1 interbody fusion with the AxiaLIF rod and pedicle screw fixation. \textbf{Lower:} Coronal and sagittal CT images demonstrating L5–S1 arthrodesis. Adapted with permission from Lippincott Williams and Wilkins/Wolters Kluwer Health: Tobler WD et al: Minimally invasive axial presacral L5–S1 interbody fusion: two-year clinical and radiographic outcomes. \textit{Spine} 36(20):E1296–E1301, 2011.
Complications

A number of recently published case reports have described rectal perforation in association with the AxiaLIF surgery. Botolin and colleagues reported the case of a 44-year-old woman with a history of anterior and posterior spinal surgeries, pelvic inflammatory disease, and diverticulitis who underwent L5–S1 axial interbody fusion. After surgery, she presented with abdominal pain, nausea, vomiting, and melena. An abdominal CT scan after intravenous and oral contrast administration demonstrated soft-tissue fluid density with fat stranding in the presacral space. There was extraluminal rectal contrast and gas consistent with rectal perforation (Fig. 5). The patient required a diverting ileostomy with intravenous antibiotics. The authors suggested that a preoperative pelvic CT scan with rectal contrast be acquired in patients at risk for adhesion formation. Mazur and colleagues reported on a patient presenting with progressive back pain and sepsis 3 weeks after an L5–S1 AxiaLIF procedure. The patient required diverting colostomy and antibiotic treatment. She went on to develop a nonunion, which was treated with revision posterior spinal fusion and no explantation of the AxiaLIF device. Siegel and colleagues reported on the development of a rectocutaneous fistula in a 35-year-old man after L5–S1 AxiaLIF. The patient presented with sepsis and bloody drainage from the surgical site. A CT fistulagram and flexible sigmoidoscopy showed evidence of a rectocutaneous fistula, which was managed with intravenous antibiotics. At 6 months, the patient developed a pseudarthrosis that required surgical revision.

Lindley and colleagues retrospectively reviewed complications in 68 patients who underwent AxiaLIF. Sixteen patients (23.5%) suffered complications including rectal perforation (2 patients), pseudarthrosis (6 patients), superficial infection (4 patients), pelvic hematoma (2 patients), and sacral fracture (2 patients). With regard to...

Pseudarthrosis is a concern when using the 2-level AxiaLIF procedure. Issack and Boachie-Adjei reported two pseudarthroses in their series: one at L4–5 and the other at L5–S1. One of these patients had a near 10-cm positive sagittal balance after AxiaLIF. Although the numbers are too small in this series to draw statistically significant conclusions, restoration of sagittal balance is likely to be important for interbody fusion. Furthermore, thorough and meticulous discectomy and endplate preparation followed by robust bone grafting are likely essential components (as in ALIF, TLIF, and PLIF) of successful arthrodesis. Hofstetter and colleagues retrospectively reviewed data obtained in 38 patients who underwent either 1-level (32 patients) or 2-level (6 patients) AxiaLIF; the mean follow-up period was 26 months. Graft subsidence eliminated partial correction of segmental lordosis. At final follow-up, 72% of L5–S1 levels exhibited bony fusion (80% of the 1-level and 33% of the 2-level AxiaLIF procedures). None of the L4–5 levels in 2-level AxiaLIF fused. Five patients required revision fusion. The authors suggested that the axial rod provides inadequate long-term anterior column support. Marchi and colleagues prospectively followed up 27 patients who underwent the AxiaLIF procedure at L4–5 and L5–S1. In minimum follow-up period of 2 years, the authors observed multiple implant-related complications including screw breakage (15%), proximal/distal transsacral rod detachment (11%), and superior rod migration (24%). At 2-year follow-up, the disc space was diminished in comparison with the preoperative status, segmental lordosis was diminished, and only 22% of all treated levels exhibited a solid fusion.

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superficial infection, we found that making a transverse incision, instead of a longitudinal incision, reduced this risk. To prevent pelvic hematoma, we recommend that all patients undergo preoperative MRI to document a clear fat plane separating the visceral peritoneum and rectum from the anterior sacral wall. If extensive vasculature is seen in this plane, AxiaLIF is contraindicated, as instruments cannot be safely passed through the presacral space. To prevent sacral fracture, the trajectory of the guide pin is critical. Too anterior a path may result in fracture of the anterior sacral cortex when either instruments or the implant is inserted. A trajectory that is too posterior may result in posterior vertebral cortical penetration with possible neural element injury.

Conclusions

The axial transsacral approach to L5–S1 provides a minimally invasive approach to interbody arthrodesis at L5–S1. Biomechanical studies on the AxiaLIF implant suggest stiffness comparable to existing interbody fixation devices. Early clinical studies on the AxiaLIF demonstrate good to excellent results in terms of pain relief and fusion rates when performing L5–S1 arthrodesis to treat degenerative disc disease and spondylolisthesis. Short-term follow-up studies on the use of the axial interbody fusion caudal to long fusion constructs demonstrate good pain relief and fusion rates and minimal blood loss. The technique, however, is exacting and complications can be major. Emerging reports demonstrate a poor fusion rate with the 2-level construct. Long-term follow-up studies to assess clinical outcome, fusion, and complications are required before axial bone graft is recommended as a standard, routine approach to achieve L5–S1 interbody arthrodesis in adult scoliosis.

Disclosure

Dr. Boachie-Adjei reports being a consultant for Baxano, K2M, DePuy, and Medtronic.

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P. S. Issack, S. Y. Kotwal, and O. Boachie-Adjei
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